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* * * * * RECONNECTED TO STN INTERNATIONAL * * * * * * SESSION RESUMED IN FILE 'HCAPLUS' AT 11:53:10 ON 29 AUG 2006 FILE 'HCAPLUS' ENTERED AT 11:53:10 ON 29 AUG 2006 COPYRIGHT (C) 2006 AMERICAN CHEMICAL SOCIETY (ACS)

COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION FULL ESTIMATED COST 2.53 169.68

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SINCE FILE TOTAL
ENTRY SESSION
FULL ESTIMATED COST
2.53 169.68

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STRUCTURE FILE UPDATES: 28 AUG 2006 HIGHEST RN 904961-01-9 DICTIONARY FILE UPDATES: 28 AUG 2006 HIGHEST RN 904961-01-9

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http://www.cas.org/ONLINE/UG/regprops.html

Uploading C:\Program Files\Stnexp\Queries\10718879a.str



chain nodes :

Young, Shawquia, Page 1

10 11 12 13

ring nodes :

1 2 3 4 5 6 7 8 9

chain bonds :

8-10 10-11 11-12 12-13

ring bonds :

1-2 1-6 2-3 3-4 4-5 5-6 5-7 6-9 7-8 8-9

exact/norm bonds :

5-7 6-9 7-8 8-9 8-10 10-11

exact bonds : 11-12 12-13

normalized bonds :

1-2 1-6 2-3 3-4 4-5 5-6

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:CLASS 11:CLASS 12:CLASS 13:CLASS

L5 STRUCTURE UPLOADED

=> d 15

L5 HAS NO ANSWERS

L5 STR

Structure attributes must be viewed using STN Express query preparation.

=> s 15

SAMPLE SEARCH INITIATED 11:53:45 FILE 'REGISTRY'

SAMPLE SCREEN SEARCH COMPLETED - 61 TO ITERATE

100.0% PROCESSED 61 ITERATIONS 3 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

72 ANSWERS

PROJECTED ITERATIONS: 752 TO 1688

PROJECTED ANSWERS: 3 TO 163

L6 3 SEA SSS SAM L5

=> s 15 full

FULL SEARCH INITIATED 11:53:48 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 1314 TO ITERATE

100.0% PROCESSED 1314 ITERATIONS

SEARCH TIME: 00.00.01

Young, Shawquia, Page 2

L7 72 SEA SSS FUL L5

=> file hcaplus
COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 166.94 336.62

FULL ESTIMATED COST

FILE 'HCAPLUS' ENTERED AT 11:53:53 ON 29 AUG 2006
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FILE COVERS 1907 - 29 Aug 2006 VOL 145 ISS 10 FILE LAST UPDATED: 28 Aug 2006 (20060828/ED)

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This file contains CAS Registry Numbers for easy and accurate substance identification.

=> s 17 L8 19 L7

=> d ed abs ibib hitstr 1-19

ANSWER 1 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN Entered STN: 28 Oct 2005

AB The title compds. I [A = alkyl, cycloalkyl, aryl, etc.; UV = R2CH:CH, R2CH:CH (wherein R2 = H, alkyl, alkoxy); W = S, O, NR3 (R3 = H, alkyl); X

(CH2) nCH2O(CH2) nCH2C.tplbond.C, NR1(CH2) nCH2C.tplbond.C, (CH2) nCH2C.tplbond.C (n = 0-3; R1 = H, alkyl, with the proviso that when

- (CH2)nCH2C.tplbond.C, UV can not be R2CHCH, unless A = acyloxy); B = aryl, heterocyclyl; Y = (CH2)n (n = 0-3); R = H, alkyl, cycloalkyl, aryl, heterocyclyl] which have PPAR agenist activity and hence can be used as antidiabetic compds, were prepared General procedures for synthesis of I such as 5-(4-(3-acetoxyprop-1-ynyl)benzylidenelthiazolidine-2,4-dione, were given (no specific synthetic example). Compds I can be used for

treatment of diabetes and diabetes-associated complications, for the treatment of diseases and conditions in which insulin resistance is the central pathophysiol. mechanism, for the treatment of diseases or conditions such as Type II diabetes, dyslipidemia, hypertension, coronery heart disease, cardiovascular disease, atherosclerosis, diabetes nephropathy, glomerulonephritis, glomerularsclerosis, nephrotic syndrome, hypertensive nephrocelerosis, polycystic ovarian syndrome, eating disorders, psoriasis, obesity, for improving cognitive functions in dementia and as aldose reductase inhibitors. Processes for the

diabetes mellitus and the diseases and conditions mediated through insulin

resistance are claimed. SSION NUMBER: 200

2005:1154538 HCAPLUS

DOCUMENT NUMBER: TITLE:

143:422344
Preparation of alkynyl substituted thiazolidinediones as antidiabetic egents
Salman, Mohammad; Sattigeri, Jitendra; Vir, Dharam;
Gangan, Vija Dattatraya
Ranbaxy Laboratories Limited, India
PCT Int. Appl., 33 pp.
CODEN: PIXXD2
Patent
Explish

INVENTOR (S):

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE:

English

PAMILY ACC. NUM. COUNT: PATENT INFORMATION:

ANSWER 2 OF 19 HCAPLUS COPYRIGHT 2006 ACS ON STN Entered STN: 28 Oct 2005

Title compds. I {A = alkyl, alkenyl, acyl, etc.; X = no atoms, O, -(CH2)nO-(CH2)n-CH2-CHCH, etc.; n = 0-3; B = aryl or heterocycle; Y = (CH2)m; m = 1-3; R1 = OR3 or NR3R4; R3 and R4 = H or alkyl; R2 = alkyl, cycloalkyl, aryl, etc.] and their pharmaceutically acceptable salts, ary prepared and disclosed as peroxisome proliferator activated receptor

agonists. Thus, e.g., II was prepared by benzylation Et ethoxy acetate

agonists. Thus, e.g., II was prepared by benzylation Et ethoxy acetate with

4-bromobenzylbromide followed by coupling with the resp. propargylic heterocycle. The activity of I was evaluated in binding assays for PPARa, PPARB and PPARy using CARLA assays and it was revealed that selected compds. of the invention displayed EC50 values in the range of 0.04 to 30 µM for PPARa, 0.03 up to 30 µM for PPARB, I as agonist of PPAR should prove useful in the PPARB, I as agonist of PPAR should prove useful in the treatment of diseases such as but not limited to type II disbetes, cardiovascular disease and obesity. Order to type II disbetes, cardiovascular disease and obesity. Pharmaceutical compas. comprising I are disclosed.

ACCESSION NUMBER: 2005:1154526 HCAPLUS

ACCESSION NUMBER: 143:440407

TITLE: Preparation of substituted phenyl propanoic acids and esters as peroxisome proliferator activated receptor (PPAR) agonists

Salam, Nohammad; Sattigeri, Jitendra A.

Ranbaxy Laboratories Limited, India PCT Int. Appl., 30 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: PT Int. Appl., 30 pp.

CODEN: PIXXD2

PATENT INFORMATION: Patent

INFORMATION: PATENT INFORMATION:

PAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE A1 WO 2005100318 20051027 WO 2005-IB1002 20050414

Young, Shawquia, Page 4

L8 ANSWER 1 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) PATENT NO. KIND DATE APPLICATION NO. DATE WO 2005100331 WO 2005100331 A2 A3 20051027 WO 2005-IB998 20050414 100311 A3 20060406
AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KP, KR, KZ, KL, LK, LK, LT, LU, LV, MA, MD, MG, MK, MM, MM, MZ, MZ, NJ, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SM, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
BM, GM, GM, KE, LS, MM, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, PR, GB, GR, HU, IE, IS, IT, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BP, BJ, CP, CG, CI, CH, GA, GN, GQ, GN, MR, MR, NE, SN, TD, TG 20060406 PRIORITY APPLN. INFO.: US 2004-562009P P 20040414 R SOURCE(S):

MARPAT 143:422344

868362-66-9P, 5-[4-[3-(N-(2-Benzothiazolyl)-N-methylamino)prop-1ynyl]benzylidenelthiazolidine-2,4-dione 868363-96-6P,

5-[4-[3-(N-(2-Benzothiazolyl)-N-methylamino)prop-1ynyl]benzyl]thiazolidine-2,4-dione
RI: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
(Usea) OTHER SOURCE(S):

(vsea) (preparation of thiazolidinediones as antidiabetic agents) 868382-68-9 HCAPLUS (2-4-7hiazolidinedione, 5-[[4-[3-(2-benzothiazolylmethylamino]-1-propynyl]phenyl]methylene]- (9CI) (CA INDEX NAME)

868363-96-6 MCAPLUS
2,4-Thiazolidinedione, 5-[[4-[3-(2-benzothiazolylmethylamino]-1-propynyl]phenyl]methyl]- [9CI] (CA INDEX NAME)

ANSWER 2 OF 19 HCAPLUS COPYRIGHT 2006 ACS ON STN (CONTINUED)

N: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, EW, BY, BZ, CA, CH,
CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, BE, ES, F1, GB, GD,
GE, GH, GM, HR, HU, ID, IL, IN, 1S, JP, KE, KO, KM, KP, KR, KZ,
LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, NN, MM, MX, MZ, NA,
NI, NO, NZ, OM, FG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL,
SM, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA,
ZM, ZW

RN: EM, GM, GM, KE, LS, MM, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZM,
AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK,
EE, ES, F1, FR, GB, GR, HU, IE, IS, IT, LT, LU, MC, NL, PL, PT,
RO, SE, S1, SK, TR, BP, BJ, CF, CG, CI, CM, GA, GN, GG, GW, MI,
NE, NE, SN, TD, TG

ORITY APPLIA IMFO: MR, NE, S PRIORITY APPLN. INFO.: US 2004-562085P P 20040414

OTHER SOURCE(S): MARPAT 143:440047

REGIOE2-31-9P
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

(preparation of substituted Ph propanoic acids and esters as peroxisome

xisome
 proliferator activated receptor (PPAR) agonists)
868082-31-9 HCAPLUS
Benzenepropanoic acid, 4-{3-{a-benzothiazolylmethylamino}-1-propynyl}α-ethoxy-, ethyl ester (9CI) (CA INDEX NAME)

3

REPERENCE COUNT:

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

ANSWER 3 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN Entered STN: 30 Jun 2005

Title compds. I (A = alkyl, alkenyl, alkynyl, etc.; X = -(CH2)n-(CH2)-0-(CH2)n, NR(CH2)n-, -(CH2)n-; n = 0-3; R = H, alkyl; Y (CH2)n; U-V = R3C-C, R3CH-CH; R3 = H, alkyl, alkoxy; W = S, O, NR4; R4 H, alkyl; R1 = alkyl, cycloalkyl, OH, etc.; R2 = H, alkyl, aryl, etc.]

and
their pharmaceutically acceptable salts, are prepared and disclosed as
agonists of PPAR receptors. Thus, e.g., II was prepared by a multi-step
process. The activity of I was evaluated in a functional and binding
assay for PPARA/6/y and it was revealed that compds. of
the invention displayed EC50 values for PPARA from 0.02 µM to
greater than 30 µM. I as an agonist of PPAR receptors should prove
useful in the treatment of diabetes. Pharmaceutical compns. comprising I
accession NUMBER: 2005:567158 HCAPLUS
DOCUMENT NUMBER: 143-93158

DOCUMENT NUMBER: TITLE:

INVENTOR (S)

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: PAMILY ACC. NUM. COUNT:

2005:567158 HCAPLUS
143:97157
Preparation of phenyl acetylene derivatives as agonists of PPAR receptors
Sattigeri, Jitendra A.; Salman, Mohammad
Ranbaxy Laboratories Limited, India
PCT Int. Appl., 44 pp.
CODEN: PIXKD2
Patent
English
1

ANSWER 3 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

$$\begin{array}{c} \text{Me} \\ \text{N} \\ \text{N} \\ \text{CH}_2 \\ \text{CH}_2 \\ \text{CH}_2 \\ \text{CH}_2 \\ \text{O} \\ \text{N} \\ \text{H} \\ \end{array}$$

L8 ANSWER 3 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN PATENT INFORMATION:

PAT	ENT .	NO.			KIN	D	DATE			APPL	CAT	ION :	NO.		D.	ATE	
						-									-		
WO .	2005	0588	13		A2		2005	0630	,	NO 2	004-	I B 4 1	43		2	0041	215
NO.	2005	0588	13		A3		2005	0825									
	W:	AE,	AG,	AL,	AM,	AT,	ΑU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA.	CH.
		CN,	co,	CR,	cu,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES.	FI,	GB.	GD.
		GE,	GH,	GM,	HR,	ΗU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC.
		LK,	LR,	LS,	LT,	w,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NA,	NI,
		NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	sc.	SD,	SE,	SG,	SK,	SL,	SY,
		TJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW
	RW:	BW,	GH,	GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,
		AZ,	BY,	KG,	ΚZ,	MD,	RU,	TJ,	TM,	ΑT,	BE,	BG,	CH,	CY,	C2,	DE,	DK,
		ĒE,	ES,	ΡI,	FR,	GB,	GR,	HU,	IE,	IS,	IT,	LT,	LU,	MC,	NL,	PL,	PT,
		RO,	SE,	SI,	SK,	TR,	BP,	BJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	G₩,	ML,
		MR,	NE,	SN,	TD,	TG											
RIORITY	APP	LN.	INFO	. :					1	JS 2	003-	5303	34 P	1	P 2	0031	217

856256-35-4P RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(Uses)

(preparation of Ph acetylene derivs. as agonists of PPAR receptors)

RN 856256-35-4 HCAPLUS

CN 2,4-Thiazolidimedione,
5-[[3-[3-(2-benzothiazolylmethylamino)-1-propynyl]4-methoxyphenyl]methyl]- (9CI) (CA INDEX NAME)

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ANSWER 4 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN Entered STN: 10 Sep 2004
```

$$F_3$$
co $\stackrel{S}{\underset{R^4}{\longrightarrow}} \stackrel{R^1}{\underset{R^2}{\longrightarrow}} R^3$

AB Title compds. [I; Rl = null, H, alkyl, alkynyl, aminoalkyl, hydroxyalkyl, (CH2)yS(CH2)xMe, etc.; x = 0-5; y = 1-5; x+y <6; R2, R3 = H, alkyl; R4 = null, H, alkyl, alkynyl, (CH2)yS(CH2)xMe, aminoalkyl, hydroxyalkyl, etc.; ≥1 of R1, R4 is present; dotted line = bond between 1 of the N atoms and the intervening C atom), were prepared Thus, 2-chloro-6-trifluoromethoxybenzothiazole (preparation given) and N-methylproparyylamine were stirred overnight to give methylprop-2-ynyl (6-trifluoromethoxybenzothiazol-2-yl)amine. The latter as the hydrochloride at 10 MM in MPP trested PC-12 cells reduced LDH release from 49.7% to 10.9% of total.

ACCESSION NUMBER: 2004:739978 HCAPLINE

ACCESSION NUMBER: 2004:739979 HCAPLUS
DOCUMENT NUMBER: 141:241548
Preparation of
trifluoromethoxypropargyleminobenzothia
RIVENTOR(S): Sterling, Jeffrey: Hayardeny, Liat; Palb, Eliezer;
Herzig, Yaacov, Lerner, David
Israel
SURCE: U.S. Pat. Appl. Publ., 25 pp.
DOCUMENT TYPE: Paking P ACCESSION NUMBER 2004:739979 HCAPLUS

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

APPLICATION NO. PATENT NO. KIND DATE DATE US 2004176430 PRIORITY APPLN. INFO.: US 2003-718879 US 2002-428093P A1 20040909 20031120 P 20021121

OTHER SOURCE(S):

R SOURCE(S): CASREACT 141:243548; MARPAT 141:243548
702659-90-3P 702659-91-4P 702659-92-5P
702659-93-6P 702659-91-7P 702659-98-1P
702659-96-9P 702659-97-0P 702659-98-1P
702659-99-2P 702660-00-2P 702660-01-3P
702660-02-4P 702660-03-5P 702660-04-6P
702660-05-7P 702660-06-8P 702660-07-9P
702660-08-0P 702660-09-1P 702660-11-5P
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of trifluoromethoxypropargylaminobenzothiazoles for (Preparation of neurol. disorders)
of neurol. disorders)
RN 702659-90-3 HCAPLUS
CN Ethanone, 1-(4-methylphenyl)-2-[2-propynyl][6-(trifluoromethoxy)-2-

L8 ANSWER 4 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) benzothiazolyl]amino] - (9CI) (CA INDEX NAME)

RN 702659-91-4 HCAPLUS CN 2-Benzothiazolamine, N,N-di-2-propynyl-6-(trifluoromethoxy)- (9CI) (CA INDEX NAME)

RN 702659-92-5 HCAPLUS
CN 2-Benzothiazolamine, N-[2-(methylthio)ethyl]-N-2-propynyl-6-(trifluoromethoxy)- (9Cl) (CA INDEX NAME)

RN 702659-93-6 HCAPLUS
CN 2-Propyn-1-amine, N-{3-[2-(methylthio)ethyl]-6-(trifluoromethoxy)-2(3H)-benzothiazolylidene]- (9CI) (CA INDEX NAME)

RN 702659-94-7 HCAPLUS
CN 2-Propyn-1-amine, N-[3-(2-propynyl)-6-(trifluoromethoxy)-2(3H)-benzothiazolylidene]- (9CL) (CA INDEX NAME)

L8 ANSWER 4 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

RN 702659-99-2 HCAPLUS
CN 2-Benzothiazolamine, N-2-propynyl-6-(trifluoromethoxy)-,
monohydrochloride
(9CI) (CA INDEX NAME)

• HC1

RN 702660-00-2 HCAPLUS CN 2-Benzothiazolamine, N-2-butynyl-6-(trifluoromethoxy)- (9CI) (CA INDEX NAME)

RN 702660-01-3 HCAPLUS
CN 2-Benzothiazolamine, N-2-butynyl-6-(trifluoromethoxy)-, monohydrochloride
(9C1) (CA INDEX NAME)

• HCl

RN 702660-02-4 HCAPLUS
CN 2-Benzothiezolamine,
N-methyl-N-(1-methyl-2-propynyl)-6-(trifluoromethoxy)(9CI) (CA INDEX NAME)

L8 ANSWER 4 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

RN 702659-95-8 HCAPLUS Ethanone, 1-(4-methylphenyl)-2-(2-(2-propynylimino)-6-(trifluoromethoxy)-3(2H)-benzothiazolyl)- (9Cl) (CA INDEX NAME)

RN 702659-96-9 HCAPLUS
CN 2-Benzothiezolamine, N-methyl-N-2-propynyl-6-(trifluoromethoxy)- (9CI)
(CA INDEX NAME)

RN 702659-97-0 HCAPLUS
CN 2-Benzothiazolamine, N-methyl-N-2-propynyl-6-(trifluoromethoxy)-,
monohydrochloride (SCI) (CA INDEX NAME)

HC1

RN 702659-98-1 HCAPLUS
CN 2-Benzothiazolamine, N-2-propynyl-6-(trifluoromethoxy)- (9CI) (CA INDEX NAME)

L8 ANSWER 4 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

RN 702660-03-5 HCAPLUS
CN 2-Benzothiazolamine,
N-methyl-N-(1-methyl-2-propynyl)-6-(trifluoromethoxy), monohydrochloride (9CI) (CA INDEX NAME)

• HCl

RN 702660-04-6 HCAPLUS
CN 2-Benzothiazolamine, N-methyl-N-2-propynyl-4-(trifluoromethoxy)- (9CI)
(CA INDEX NAME)

RN 702660-05-7 HCAPLUS
CN 2-Benzothiazolamine, N-methyl-N-2-propynyl-4-(trifluoromethoxy)-,
monohydrochloride (9CI) (CA INDEX NAME)

ANSWER 4 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

● HCl

702660-06-8 HCAPLUS
2-Propyn-1-amine, N-[3-methyl-6-(trifluoromethoxy)-2(3H)-benzothiazolylidene)- (9CI) (CA INDEX NAME)

702660-07-9 HCAPLUS
2-Propyn-1-amine, N-{3-methyl-6-(trifluoromethoxy)-2(3H)-benzothiazolylidene}-, monohydrochloride (9CI) (CA INDEX NAME)

• HC1

702660-08-0 HCAPLUS

nethyl-2-(methyl-2-propynylamino)-6-(trifluoromethoxy), iodide (9CI) (CA INDEX NAME)

L8 ANSWER 4 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN

(Continued)

702660-09-1 HCAPLUS
2-Propyn-1-amine, N-[3-methyl-4-(trifluoromethoxy)-2(3H)-benzothiazolylidene]- (9CI) (CA INDEX NAME)

702660-11-5 HCAPLUS 2-Propyn-1-amine, N-[3-methyl-6-(trifluoromethoxy)-2(3H)-benzothiazolylidene]-, monohydriodide (9CI) (CA INDEX NAME)

• ні

ANSWER 5 OP 19 HCAPLUS COPYRIGHT 2006 ACS ON STN Entered STN: 10 Jun 2004

The invention provides title compds. I [wherein R1 is present or absent, and when present = H, C1-6 alkyl, C1-6 alkynyl, (CH2)yS(CH2)xCH3, C1-6 aminoalkyl, C1-6 hydroxyalkyl, or (CH2)nC0(C6H4)(CH2)R2; R2 = H or C1-4 alkyl; R3 = H or C1-4 alkyl; R4 is present or absent, and when present = H, C1-6 alkyl, C1-6 alkynyl, (CH2)yS(CH2)xCH3, C1-6 aminoalkyl, C1-6 hydroxyalkyl, or (CH2)nCO(C6H4)(CH2)R2; N = 1-6; wherein x = 0.5; y =

such that (x+y) < 6; at least one of R1 or R4 is present; dashed line = bond between one of two N atoms and the intervening C atom; and the

ound is charged when both R1 and R4 are present; including any specific enantiomer, or any pharmaceutically acceptable salt]. The invention also provides a method for treating a neurol. disorder or multiple sclerosis

administering a therapeutically effective amount of any of the compds. I. Neurol. disorders listed in claims include Parkinson's disease, Alzheimer's disease, amyotrophic lateral sclerosis, stroke, neuromuscular disorders, schizophrenia, cerebral infarction, head traums, glaucoma, facialis, and Huntington's disease. The use of I for destroying or inhibiting the proliferation of microbes or fungi is also claimed. For instance, hydrazinolysis of 6-trifluoromethoxy-2-aminobenzothiazole with NHANNZ.H2SO4 and NHANNZ.H2SO4 and HANNZ-H2SO4 H2SO4 H

with SOC12 (neat) at 65° to give 2-chloro-6trifluoromethoxybenzothiazole. This chloride (crude) was treated with
N-methylpropargylamine to give invention compound II, also isolated as
II.RCI (III) by precipitation from EroH/HCl using Et20. III showed
neuroprotective activity against MPP+ toxicity, both in vitro (PC-12
cells) and in vivo (aice). At 10 mg/kg, twice daily, II gave complete
protection of mice against mortality in an exptl. allergic
encephelomyelitie (EAB) model of multiple sclerosis.

ACCESSION NUMBER: 2004:470953 HCAPLUS
DOUMENT NUMBER: 141:36603
TITLE: Propargyl-trifluoromethoxy-amino-benzothiszole
derivatives with neuroprotective activity. and their

141:38630 Propargyl-trifluoromethoxy-amino-benzothiazole derivatives with neuroprotective activity, and their

Young, Shawquia, Page 7

L8 ANSMER 5 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)
preparation, pharmaceutical compositions, and use
Sterling, Jeffrey; Hayardeny, Liat; Flab, Eliezer;
Herzig, Yaacov; Lerner, David
Teva Pharmaceutical Usa, Inc.
SOURCE:

DOCUMENT TYPE:

HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)
preparation, pharmaceutical compositions, and use
Sterling, Jean-Caption, pharmaceutical usa, Inc.
PCT Int. Appl., 78 pp.
CODEN: PIXXD2
DOCUMENT TYPE:

DOCUMENT TYPE: LANGUAGE:

English

PAMILY ACC. NUM. COUNT: PATENT INFORMATION:

												LICAT						
	WO	2004	0477	56		A2		2004	0610			2003-						
	WO	2004	0477	56		A3		2004	0708									
		W:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB	, BG,	BR,	BW,	BY,	BZ,	CA,	CH,
			CN,	co,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ	, EC,	EE,	EG,	ES,	FI,	GB,	GD,
												, JP.						
			LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG	, MK,	MN,	MW.	MX,	MZ.	NI.	NO.
			NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC	, SD,	SE,	SG.	SK.	SL.	SY,	TJ,
												, VN,						
		RW:	BW,	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL	, sz,	TZ,	UG,	ZM.	ZW,	AM,	AZ,
			BY,	KG,	KZ,	MD,	RU,	TJ,	TM,	AT,	BE	, BG,	CH,	CY,	CZ,	DE,	DK,	EE,
			ES,	FI,	FR,	GB,	GR,	HU,	IE,	IT,	LU	, MC,	NL,	PT,	RO.	SE.	SI,	SK,
			TR,	BF,	BJ,	CP,	CG,	CI,	CM,	GΑ,	GN	, GQ,	GW,	ML.	MR,	NE.	SN,	TD,
'G																		
												2003-						
												2003-						
												2003-						
	ΕP	1569	641			A2		2005	0907		EP.	2003-	7871	12		2	0031	120
		R:	AT,	BE,	CH,	DE,	DK,	ES,	PR,	GB,	GR	, IT,	LI,	LU,	NL,	SE,	MC,	PT,
			IE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	λL	, TR,	BG,	CZ,	EE,	HU,	SK	
	CN	1741	802			A		2006	0301		CN :	2003-	8010	9082		2	0031	120
	JΡ	2006	50739	50		T2		2006	0302		JP :	2004-	5557	02		2	0031	120
	NO	2005	0029	79		Α		2005	0819	1	NO :	2005-	2979			2	0050	617
RIOF	RITY	APP	LN.	INPO	. :					1	US :	2004 - 2005 - 2002 -	3015	40		1 2	0021	121
										1	WO:	2003 -	US37	592		4 2	0031	120

WO 2003-US37592 W 20031120

OTHER SOURCE(S): CASREACT 141:38603; MARPAT 141:38603

IT 703659-98-1P, Prop-2-ynyl(6-trifluoromethoxybenzothiazol-2-y1)amine 702660-06-8P, (3-Methyl-6-trifluoromethoxy-3H-benzothiazol-2-ylidene)(prop-2-ynyl)amine
RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses) (drug candidate; preparation of (propargylamino)(trifluoromethoxy)benzothiaz ole derivs. as neuroprotectants)
RN 702659-98-1 HCAPLUS
NAME)

ANSWER 5 OF 19 HCAPLUS COPYRIGHT 2006 ACS OR STN (Continued)

NH-CH1-C=CH

702660-06-8 HCAPLUS
2-Propyn-1-amine, N-(3-methyl-6-(trifluoromethoxy)-2(3H)-benzothiazolylidene]- (9CI) (CA INDEX NAME)

He

1T 702659-90-3P, 2-[Prop-2-ynyl(6-trifluoromethoxybenzothiazol-2-yl)laminol-1-p-tolylethanone 702659-91-4P, Diprop-2-ynyl(6-trifluoromethoxybenzothiazol-2-ylamine 702659-92-5P, (2-Methylaulfanylethyl) [prop-2-ynyl) (6-trifluoromethoxybenzothiazol-2-ynyl) [6-trifluoromethoxybenzothiazol-2-ynyl) [6-trifluoromethoxybenzothiazol-2-ynyl) [6-trifluoromethoxybenzothiazol-2-ynyl) [6-trifluoromethoxy-3H-benzothiazol-2-ynyl] [6-trifluoromethoxy-3H-benzothiazol-2-ynyl] [6-trifluoromethoxy-3H-benzothiazol-2-ynyl] [6-trifluoromethoxyl) benzothiazol-3-ynyl] [6-trifluoromethoxyl) benzothiazol-3-ynyl] [6-trifluoromethoxybenzothiazol-2-ynyl] [6-triflu

ANSWER 5 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) /N-CH2-C=CH F₃C

702659-95-8 HCAPLUS
Ethanone, 1-(4-methylphenyl)-2-[2-(2-propynylimino)-6-(trifluoromethoxy)-3(2H)-benzothiazolyl)- (9CI) (CA INDEX NAME)

702659-96-9 HCAPLUS 2-BenZothiazolamine, N-methyl-N-2-propynyl-6-(trifluoromethoxy)- (9CI) (CA INDEX NAME)

702659-97-0 HCAPLUS
2-Benzothiazolamine, N-methyl-N-2-propynyl-6-(trifluoromethoxy)-,
monohydrochloride (9CI) (CA INDEX NAME)

● RC1

702659-99-2 HCAPLUS
2-Benzothiazolamine, N-2-propynyl-6-(trifluoromethoxy)-, hydrochloride (9CI) (CA INDEX NAME)

ANSWER 5 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) Ethanone, 1-(4-methylphenyl)-2-[2-propynyl16-(trifluoromethoxy)-2-benzothizaclyl]amino|-(9C1) (CA NIDEX NAME)

702659-91-4 HCAPLUS 2-BenZothiezolamine, N,N-di-2-propynyl-6-(trifluoromethoxy)- (9CI) (CA INDEX NAME)

702659-92-5 HCAPLUS
2-Benzothiazolamine, N-[2-{methylthio}ethyl]-N-2-propynyl-6-{trifluoromethoxy}- (9CI) (CA INDEX NAME)

702659-93-6 HCAPLUS
2-Propyn-1-amine, N-[3-[2-(methylthio)ethyl]-6-(trifluoromethoxy)-2(3H)-benzothiazolylidene)- (9CI) (CA INDEX NAME)

702659-94-7 HCAPLUS
2-Propyn-1-amine, N-{3-(2-propynyl)-6-(trifluoromethoxy}-2(3H)-benzothiazolylidene)- (9CI) (CA INDEX NAME)

L8 ANSWER 5 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

702660-00-2 HCAPLUS 2-Benzothiazolamine, N-2-butynyl-6-(trifluoromethoxy)- (9CI) (CA INDEX

702660-01-3 HCAPLUS 2-Benzothiazolamine, N-2-butynyl-6-(trifluoromethoxy)-, monohydrochloride (9CI) (CA INDEX NAME)

RN 702660-02-4 HCAPLUS
CN 2-Benzothiezolamine,
N-methyl-N-(1-methyl-2-propymyl)-6-(trifluoromethoxy)(9CI) (CA INDEX NAME)

RN 702660-03-5 HCAPLUS
CN 2-Benzothiazolamine,
N-methyl-N-(1-methyl-2-propynyl)-6-(trifluoromethoxy), monohydrochloride (9CI) (CA INDEX NAME)

ANSWER 5 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

● HC1

702660-04-6 HCAPLUS 2-Benzothiazolamine, N-methyl-N-2-propynyl-4-(trifluoromethoxy)- (9CI) (CA INDEX NAME)

702660-05-7 HCAPLUS
2-Benzothiazolamine, N-methyl-N-2-propynyl-4-(trifluoromethoxy)-monohydrochloride (9CI) (CA INDEX NAME)

702660-07-9 HCAPLUS
2-Propyn-1-emine, N-[3-methyl-6-(trifluoromethoxy)-2(3H)-benzothiszolylidenej-, monohydrochloride (9CI) (CA INDEX NAME)

ANSWER 5 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

• ні

L8 ANSWER 5 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN

• HCl

RN 702660-08-0 HCAPLUS
CN Benzothiazolium,
3-methyl-2-(methyl-2-propynylamino)-6-(trifluoromethoxy), iodide (9C1) (CA INDEX NAME)

• 1

702660-09-1 HCAPLUS
2-Propyn-1-amine, N-[3-methyl-4-{trifluoromethoxy}-2(3H)-benzothiezolylidene]- (9CI) (CA INDEX NAME)

702660-11-5 HCAPLUS
2-Propyn-1-amine, N-[3-methyl-6-(trifluoromethoxy)-2(3H)-benzothiazolylidene]-, monohydriodide (9CI) (CA INDEX NAME)

ANSWER 6 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN Entered STN: 06 Oct 2000

The photog. film contains at least 1 kind of compound selected from R11CCO(CH2)mCO2R12, R21OCO(CH21)CO2R22, R31OCO(CH21)CO2R23, R41R42R43COH, and X-1(CH2)q-O(CO)R51)r [R11, R12, R21, R22 = C4-10-alkyl; m,n = 2-10; R31, R32 = C3-24-alkyl; p = 2-10; R41 = alkyl, alkenyl; R42, R43 = H, alkyl, alkenyl; X = 5- to 7-membered saturated hydrocarbon AB

ring, q = 0-2; r = 1-3; R51 = C4-16-alky], and at least 1 radical scavenger selected

from Xa1-(C(Ra1):Y)n-Xa2 [Xa1, Xa2 = -ORa3, -N(Ra4)Ra5; Ra3 = H, group capable of becoming H upon hydrolysis; Ra4, Ra5 = H, alkyl, alkenyl,

, heterocycle, sulfonyl, acyl, etc.; Y = C(Ra2), N; Ra1, Ra2 = H, substituent; n 20), I (Ra6-10 = H, alkyl, alkenyl, aryl, etc.], II [Ra15 = H, alkaline metal, quaternary ammonium; Ra16, Ra17 = H, halo,

alkyl,
aryl, etc.; Xa = O, substituted iminol, and RaisRa2NORais [Rais = alkyl,
alkenyl, aryl, heterocycle, acyl, sulfonyl; Rais = alkyl, alkenyl, aryl,
etc.; Ra20 = H, alkyl, alkenyl, aryl, etc.].
ACCESSION NUMBER:
DOCUMENT NUMBER:
130:288786
Silver halide color photographic film with excellent
shelf life, reduced fog, and high sensitivity
Kawabe, Satomi; Hoshino, Hiroyuki
Konica Co., Japan
Jpn. Kokai Tokkyo Koho, 67 pp.
CODEN: JIXXAAP
PAHLY ACC. NUM. COUNT:
1 alkyl.

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 2000275802	A2	20001006	JP 1999-79969	19990324
PRIORITY APPLN. INFO.:			JP 1999-79969	19990324

OTHER SOURCE(S): MARPAT 131:288786

IT 161765-65-7

RL DEV (Device component use); USES (Uses)

(in Ag halide color photog. film with excellent shelf life, reduced fog, and high sensitivity)

RN 161765-65-7 RCAPLUS

CN 2-Benzothiazolamine, N-2-butynyl- (9CI) (CA INDEX NAME)

LB ANSWER 6 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

L8 ANSWER 7 OF 19 HCAPLUS COPYRIGHT 2006 ACS ON STN R: CH, DE, FR, GB, L1 19 2001511140 T2 20010807 JP 1998-5325 PRIORITY APPLN. INFO: DE 1997-1970 JP 1998-532559 DE 1997-19704134 19980204 A 19970204 WO 1998-EP581 A 19980204 OTHER SOURCE(S): MARPAT 129:148990
IT 210834-74-5P 210834-81-4P
RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of substituted
2-(2,4(1H,3H)-pyrimidindion-3-yl)benzthiazoles
as herbicides, desiccants, and defoliants)
RN 210834-74-5 HAZPLUS
CN 2,4(1H,3H)-Pyrimidinedione, 3-[4-chloro-6-fluoro-2-(methyl-2-propynylamino)-7-benzothiazolyl)-1-methyl-6-(trifluoromethyl)- (9CI) (CA INDEX NAME)

(Continued)

210834-81-4 HCAPLUS
2.4(1H,3H)-Pyrtmidinedione, 3-[4-chloro-6-fluoro-2-(methyl-2-propynylamino)-7-benzothiazolyl]-6-(trifluoromethyl)- (9CI) (CA INDEX NAME)

210834-92-7P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

Young, Shawquia, Page 10

ANSWER 7 OF 19 HCAPLUS COPYRIGHT 2006 ACS ON STN Entered STN: 26 Aug 1998

AB Title compds. (I; R1 = alkyl, haloalkyl; R2 = H, halo; R3 = halo; R4 = alkyl; R5 = alkyl, haloalkyl, cyanoalkyl, alkoxyalkyl, alkenyl, halogenalkenyl, alkynyl, cycloalkylalkyl, Ph, phenylalkyl; R4R5 = tetramethylene; R6 = H, amino, alkyl), were prepared as herbicides, desiccants, and defoliants (no data). Thus, 4-chloro-6-fluoro-2-(pyrrolidin-1-yl)-7-(6-trifluoromethyl-2,4(lH,3H)-pyrimidindion-3-yl)-8-(byenzethiazole (preparation given) in 2-butanone was treated with K2CO3 and Mel to give 4-chloro-6-fluoro-7-(1-methyl-6-trifluoromethyl-2,4(lH,3H)-pyrimidindion-3-yl)-2-(pyrrolidin-1-yl)benzothiazole.

ACCESSION NUMBER: 1998:543070 HCAPJUS
DOCUMENT NUMBER: 1998:543070 HCAPJUS
TITLE: Preparation of substituted
2-(2,4(lH,3H)-pyrimidindion-3-yl)-benzthiazoles as herbicides, desiccants, and defoliants.

INVENTOR(S): Zagar, Cyrill; Heistracher, Elisabeth; Reinhard, Robert; Hamprecht, Gerhard; Menges, Markus; Menke, Olaf; Schafer, Peter; Westphalen, Karl-Otto;

Misslitz,

Ulf; Walter, Helmut BASF Aktiengesellschaft, Germany; et al. PCT Int. Appl., 40 pp. CODEN: PIXXD2 Patent German PATENT ASSIGNEE (S) : SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

	PA'	PENT	NO.			KIN	D	DATE	:	AP	PLIC	AT:	ON	NO.		D	ATE	
							-									-		
	WO	9833	796			A1		1998	0806	WO	199	8 - E	EP58	1		1	9980	204
		W:	CA,	JP,	US													
		RW:	AT,	BE,	CH,	DE,	DK,	ES,	FI,	FR, G	B, G	R,	IE.	IT,	LU,	MC.	NL.	PT.
SE																		
	CN	1194	1645			A		1998	0930	CN	199	6-1	1966	07		1	9960	826
	CA	2279	9644			AA		1998	0806	CA	199	8-2	279	644		1	9980	204
	EP	9582	195			A1		1999	1124	EP	199	8-5	9069	10		1	9980	204

ANSWER 7 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) (prepn. of substituted 2-(2,4(1H,3H)-pyrimidindion-3-yl)benzthiazoles as herbicides, desiccants, and defoliants) 210834-92-7 HCAPLUS 2,7-Benzothiazolediamine, 4-chloro-6-fluoro-N2-methyl-N2-2-propynyl-

(9CI) (CA INDEX NAME)

REFERENCE COUNT: THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

ANSWER 8 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN Entered STN: 10 Jul 1998

Title compds. [I; R1 = H, slkyl, haloslkyl; R2 = cyano, alkyl, haloslkyl, alkoxy, haloslkoxy, alkylthio, haloslkylthio, elkylsulfinyl, haloslkylsulfinyl, alkylsulfinyl, alkylsulfinyl, alkylsulfinyl, alkylsulfinyl, alkylsulfinyl, alkylsulfinyl, alkylsulfinyl, alkylsulfinyl, alkylsulfinyl, haloslkyl; R4 = H, halo; R5 = H, halo, cyano, alkyl, haloslkyl, haloslkyl, panosikyl, hydroxyalkyl, alkoxyl, haloslkoxyz Z = N:C(R8iO, N:C(XR8iS; X = bond, O, S, SO, SOZ, NH, NR7; R6, R7 = alkyl, haloslkyl, cyanosikyl, hydroxyalkyl, alkenyl, cyanosikynyl, haloslkenyl, alkynyl, cyanosikynyl, haloslkenyl, alkynyl, cyanosikynyl, haloslkyl, alkenyl, cyanosikyl, haloslkyl, alkysulfinyisikyl, alkoxyarbonylsikyl, etc.; R8R7 = (CR2i3, (CR2i4, CR2i3, CR2i3), CR2i3), CR2i3, (preparation

given) was stirred with dimethyldisulfide and tert-Bu nitrite in CH2Cl2

give 4-chloro-7-(4-chloro-5-difluoromethoxy-1-methyl-1H-pyrazol-3-yl)-6-fluoro-2-(methylthio)benzothiazole. The latter at 15.6 and 31.2 g/ha postemergent showed good activity against broadleaf weeds.

ACCESSION NUMBER: 1998:424249 HCAPLUS
DOCUMENT NUMBER: 129:81727

DOCUMENT NUMBER: TITLE:

129:81737
Preparation of pyrazole-3-ylbenzazoles as herbicides, desiccants, and defoliants.
Zagar. Cyrill; Hamprecht, Gerhard; Menges, Markus; Menke, Olaf; Schafer, Peter; Westphalen, Karl-Otto; Misslitz, Ulf; Walter, Helmut
Basf A.-G., Germany
PCT Int. Appl., 81 pp.
CODEN: PIXXD2
Parent INVENTOR (S):

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE:

PAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE 7090 A2 19980625 WO 1997-EP6715 19971201 7090 A3 19980917 AL, AU, BG, BR, BY, CA, CN, CZ, GE, HU, IL, JP, KR, KZ, LT, LV, MX, NO, NZ, PL, RO, RU, SG, SI, SK, TR, UA, US, UZ, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM WO 9827090

ANSWER 8 OF 19 HCAPLUS COPYRIGHT 2006 ACS ON STN (Continued)
pyrazol-3-yl]-6-fluoro-N-methyl-N-2-propynyl- (9CI) (CA INDEX NAME)

L8 ANSWER 8 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)
RN: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IB, IT, LU, MC, NL, PT, CA 1997-2275611 AU 1998-58536 CA 2275611 19980625 AU 9858536 AU 744339 EP 944623 20020221 EP 1997-954346 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, NL, SE, PT, IE, SI, FI, RO CN 1244865 20000216 CN 1997-181435 20000216 20040414 20001024 20010522 20011031 20040915 19990615 20020201 19990813 1145626 BR 1997-13947 JP 1998-527237 IL 1997-130469 AT 1997-954346 ZA 1997-11235 TW 1997-86118977 NO 1999-2924 BR 9713947 19971201 JP 2001506641 19971201 19971201 19971201 19971215 19971216 IL 130469 IL 130469 AT 275146 ZA 9711235 TW 474916 NO 9902924 NO 313880 KR 2000057600 US 6232470 BG 63873 20021216 20000925 20010515 KR 1999-705383 US 1999-331065 19990616 19990616 BG 1999-103554 DE 1996-19652240 20030430 PRIORITY APPLN. INFO.: W 19971201 WO 1997-EP6715

OTHER SOURCE(s): MARPAT 129:81727

IT 209346-54-3P 209347-06-8P
R1: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of pyrazole-3-ylbenzazoles as herbicides, desiccents, and defoliants)
RN 209346-54-3 MCAPLUS
CN 2-Benzothiazolamine, 4-chloro-7-(4-chloro-5-(difluoromethoxy)-1-methyl-1H-pyrazol-3-yl]-6-fluoro-N-methyl-N-2-propynyl- (9CI) (CA INDEX NAME)

N 209347-06-8 HCAPLUS N 2-Benzothiazolamine, -chloro-7-[4-chloro-1-methyl-5-(trifluoromethyl)-1H-

ANSWER 9 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN Entered STN: 05 Dec 1996

A photog, element comprises a support having situated thereon a silver halide emulsion, the emulsion comprising an alkynylmmine compound of the formula! I wherein Z represents atoms necessary to complete a AB 5-10-membered

5-10-membered
heterocyclic ring system, RI represents hydrogen or alkyl of from 1 to 5
carbon atoms, and R2 represents hydrogen, alkyl, aryl, heteroaryl,
carbocyclic, or heterocyclyl and at least one dihydroxy aryl compound
represented by formula II or III wherein R3 to R12 are independently
selected from the group consisting of hydrogen, hydroxy, sulfonate, and
alkyl of from 1 to 5 carbon atoms and wherein at least two of such groups
represent a hydroxy group.
ACCESSION NUMBER: 1996:713632 HCAPLUS
DOWNMENT MUMBER: 126-96802

126 96802

DOCUMENT NUMBER: TITLE:

126:96802
Photographic element and method of making silver halide emulsion
Eikenherry, Jon N.; Bernard, Robert E.
Eastman Kodak Company, USA

INVENTOR (5) :

English

PATENT ASSIGNEE (S): SOURCE: U.S., 12 pp. CODEN: USXXAM DOCUMENT TYPE:

PAMILY ACC. NUM. COUNT: PATENT INFORMATION:

LANGUAGE:

PATENT NO. KIND DATE APPLICATION NO. DATE US 5576170 JP 08339047 PRIORITY APPLN. INPO.: US 1995-430954 JP 1996-109922 US 1995-430954 A A2 19961119 19950428 19961224 A 19950428

OTHER SOURCE(S): MARPAT 126:96802

IER SOURCE(8): MARPAT 126:96802 85902-43-8 161765-65-7 161765-68-0 161765-70-4 175841-12-0 175841-13-1 RL: TEM (Technical or engineered material use); USES (Uses) (eilver halide photog. emulsions with improved sensitivity and reduced fog containing dihydroxyaryl compds. and) 85902-43-6 HCAPUIS

ANSWER 9 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (C2-Benzothiazolamine, N-2-propynyl- (9CI) (CA INDEX NAME) (Continued)

. ИН- СН2-С= СН

161765-65-7 HCAPLUS
2-Benzothiazolamine, N-2-butynyl- (9CI) (CA INDEX NAME)

NH-CH2-C= C-Me

161765-68-0 HCAPLUS
2-Benzothiazolamine, N-(3-phenyl-2-propynyl)- (9CI) (CA INDEX NAME)

NH-CH2-C= C-Ph

161765-70-4 HCAPLUS 2-Benzothiazolamine, N-2-butynyl-5-methyl- (9CI) (CA INDEX NAME)

_ NH- CH2- C== C- ме

175841-12-0 HCAPLUS 2-Benzothiazolamine, 5-methyl-N-2-propynyl- (9CI) (CA INDEX NAME)

NH-CH2-C=CH

175841-13-1 HCAPLUS
2-Benzothiazolamine, 5-chloro-N-2-propynyl- (9CI) (CA INDEX NAME)

L8 ANSWER 10 OF 19 HCAPLUS COPYRIGHT 2006 ACS ON STN ED Entered STN: 03 May 1996 AB A number of 2-(elkymylamino)-substituted heterocycles have been synthesized.

nesized.
These heterocycles rearrange in the presence of silver(I) and gold(I) salts to give novel 2M-pyrimido[2,1-b]benzoxazoles, 2M-pyrimido[2,1-b]benzoxazoles, 2M-pyrimido(2). Two of the b]benzothiazoles, and a 2M-pyrimido(2,1-b]benzoselenazole. Two of the

2H-pyrimido[2,1-b]benzoxazoles were isolated in good yield. The kinetics of the silver tetrafluoroborate-catalyzed rearrangements of selected (alkynylamino)benzoxazoles and benzothiazoles have been examined by IH

in CD3CN. Factors affecting the electron densities of the triple bond

of the nitrogen atom in the heterocycle are important in influencing the rate of rearrangement.

ACCESSION NUMBER: 1996:259706 HCAPLUS

DOCUMENT NUMBER: 125:10669

TITLE: Facile Rearrangements of Alkynylamino Heterocycles

1996:12570th HAAPLUS
125:10669
Facile Rearrangements of Alkynylamino Heterocycles
with Noble Metal Cations
Lok, Roger; Leone, Ronald E.; Williams, Antony J.
Esstman Kodak Company, Rochester, NY, 14650, USA
Journal of Organic Chemistry (1996), 61(10), 3289-97
CODEN: JOCEAH; ISSN: 0022-3263
American Chemical Society
Journal CORPORATE SOURCE: SOURCE:

PUBLI SHER :

. КИ−СИ2−С СН

161765-65-7 HCAPLUS

2-Benzothiazolamine, N-2-butynyl- (9CI) (CA INDEX NAME)

L8 ANSWER 9 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

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ANSWER 11 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN Entered STN: 22 Mar 1996
```

$$z \stackrel{\text{CN}(\mathbb{R}^1) \, C(\mathbb{R}^2) \, (\mathbb{R}^3) \, C \equiv C\mathbb{R}^4}{||}$$

A process is disclosed for preparing a photog. emulsion utilizing an alkynylamine compound as a grain growth modifier. Specifically, the

ent invention provides a process of preparing a photog, emulsion comprising introducing silver ions, halide ions, and a grain growth modifier having the structure I, wherein Z represents atoms necessary to complete a 5-9-membered heterocyclic ring system, RI, RZ and RI independently represent hydrogen or a lower alkyl of from 1 to 5 carbon atoms, and R4 represents hydrogen or an aliphatic, carbocyclic, or heterocyclic group, which may be substituted or unsubstituted, into a dispersing medium containing

silver halide seed grains and maintaining the dispersing medium

containing the seed grains and maintaining the dispersing medium containing the seed grains, silver ions, halide ions, and grain growth modifier at a pH in the range from about 4.5 to about 10 and a pAg in the range from about 6.0 to about 9.5.

ACCESSION NUMBER: 1996:169215 HCAPLUS

1996:169215 HCAPLUS DOCUMENT NUMBER:

TITLE:

1996:169215 HCAPLUS 124:30239 Process of forming a photographic emulsion Men, Xin; Lok, Roger Eastman Kodak Company, USA U.S., 20 pp. CODEN: USXXAW INVENTOR(S):

PATENT ASSIGNEE (S) :

SOURCE:

DOCUMENT TYPE: LANGUAGE:

English FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE		
US 5491056	A	19960213	US 1994-296567	19940826		
JP 08069072	A2	19960312	JP 1995-240480	19950828		
PRIORITY APPLN. INFO.:			US 1994-296567 A	19940826		

OTHER SOURCE(S): MARPAT 124:302399

R SOURCE(S): MARPAT 124:302399

85902-43-9 161765-56-7 161765-68-0

161765-70-4 161765-71-5 175841-10-0

175841-12-0 175841-13-1

RE: TEM (Technical or engineered material use); USES (Uses)

(silver halide photog. emulsion preparation using silver halide grain

growth

th modifier of) 85902-43-8 HCAPLUS 2-Benzothiazolamine, N-2-propynyl- (9CI) (CA INDEX NAME)

ANSWER 11 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

NH- CH2-C

161765-65-7 HCAPLUS 2-Benzothiazolamine, N-2-butynyl- (9CI) (CA INDEX NAME)

NH-CH2+C==C-Me

161765-68-0 HCAPLUS
2-Benzothiazolamine, N-(3-phenyl-2-propynyl)- (9CI) (CA INDEX NAME)

NH-CH2-C=C-Ph

161765-70-4 HCAPLUS
2-Benzothiazolamine, N-2-butynyl-5-methyl- (9CI) (CA INDEX NAME)

_ NH- CH2- С= С- Ме

161765-71-5 HCAPLUS 2-Benzothiazolamine, N-2-butynyl-5-chloro- (9CI) (CA INDEX NAME)

_ NTH — CH2 — C === C — Me

175841-10-8 HCAPLUS
2-Benzothiazolamine, 5-methoxy-N-2-propynyl- (9CI) (CA INDEX NAME)

L8 ANSWER 11 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

NH-CH2-C=CH

175841-12-0 HCAPLUS
2-Benzothiazolamine, 5-methyl-N-2-propynyl- (9CI) (CA INDEX NAME)

^ N NH-CH2-C=CH

175841-13-1 HCAPLUS
2-Benzothiazolamine, 5-chloro-N-2-propynyl- (9CI) (CA INDEX NAME)

, NH — CH2 — С== CH

ANSWER 12 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN Entered STN: 28 Sep 1995

$$Y^{1} \longrightarrow X \longrightarrow H \\ N - CH_{2}C = CR^{1}$$

A method of finishing an emulsion comprises providing Ag halide grains, adding to the emulsion in an amount between .apprx.0.005 and 0.10

and any to the emulsion in an amount between .apprx.0.005 and 0.10 mmol/per mole of Ag of the compound I [X = 0, S, Se; Rl = alkyl or substituted alkyl

alkyl

or aryl or substituted aryl; Y1 and Y2 individually = H, alkyl groups or
an aromatic nucleus or together = the atoms necessary to complete a
cyclic

structure containing C, O, Se, or N atoms necessary to complete a fused
aromatic

aromatic
nucleus or an alicyclic structure]. A photog, element comprising the Ag
halide emulsion is also claimed.

ACCESSION NUMBER:
1995:818680 HCAPLUS
DOCUMENT NUMBER:
123:21987
A class of compounds which increases and stabilizes
photographic speed.

INVENTOR(S):
Eikenberry, Jon Nathan; Lok, Roger; Chen, Chung Yuan
Eastman Kodak Co, USA
EUL. PATE ADPLI, 16 pp.
CODEN: EFXEDW

DOCUMENT TYPE:
Patent

Patent English

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. EP 658803 EP 658803 A1 19950621 B1 19980902 APPLICATION NO. DATE EP 1994-119840 19941215 19960319

Young, Shawquia, Page 13

ANSWER 12 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)
161765-65-7P
RL: MOA (Modifier or additive use); SPN (Synthetic preparation); PREP
(Preparation); USES (Uses)
 (compds. which increase and stabilize photog. speed.)
161765-65-7 HCAPLUS
2-Benzothiazolamine, N-2-butynyl- (9CI) (CA INDEX NAME)

NH-CH2-C=C-Me

ANSWER 13 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN
Entered STN: 14 Jun 1995
Substituted 2, 5-diaryl-4-isothiazolin-3-ones (Markush included) are
disclosed, as are their synthesis, pharmaceutical prepns. containing
, and
their use in the treatment of thrombosis and especially inflammation.
N-(4-methylphenyl)-5-phenyl-4-isothiazolin-3-one (preparation given) had

IC50 of 13.2 µM in a bovine nasal septum cartilage degradation assay.

ACCESSION NUMBER: 1995:607986 HCAPLUS

DOCUMENT NUMBER: 123:47905

TITLE: Substituted 2,5-diaryl-4-isothiazolin-3-ones as

133:47905
Substituted 2,5-diaryl-4-isothiazolin-3-ones as antiinflemmatory and antithrombotic agents
Petratite, Joseph J.; Sherk, Susan R.
The Dupont Merck Pharmaceutical Company, USA U.S.. 13 pp.
CODEN: USXXXAM

INVENTOR(S):
PATENT ASSIGNEE(S):
SOURCE:

Patent English

LANGUAGE: E PAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE US 5411977 PRIORITY APPLN. INFO.: 19950502 US 1993-40771 US 1993-40771 19930331

OTHER SOURCE(s): MARPAT 123:47905

IT 164395-92-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT

Reactant or reagenty (synthetic preparation); PREP (Preparation (Reactant or reagent) isothiazolinones as antiinflammatory and antithrombotic agents, and their preparation)
164395-92-0 HCAPLUS
2-Propymamide, N-2-benzothiazoly1-3-pheny1- (9CI) (CA INDEX NAME)

ANSWER 14 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN

167307-91-7 HCAPLUS 5-Benzothiazoleaulfonic acid, 2-[(3-phenyl-2-propynyl)amino]-, monosodium salt (9C1) (CA INDEX NAME)

● Na

167307-92-8 HCAPLUS
Phosphonic acid, [4-[(5-methyl-2-benzothiazolyl)amino]-2-butynyl]-,
monopotaseium salt (9CI) (CA INDEX NAME)

● K

167307-93-9 HCAPLUS 6-Benzothiazolecarboxylic acid, 2-(2-butynylamino)-5-chloro- (9CI) (CA INDEX NAME)

167307-94-0 HCAPLUS 5-Benzothiazolol, 2-(2-butynylamino)-, dihydrogen phosphate (ester), monopotassium salt (9CI) (CA INDEX NAME)

ANSMER 14 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN
Entered STN: 09 Jun 1995
A neg. or reversal photog. element comprises a photog. emulsion and a H2O soluble photog. sensitivity increasing alkynylamine compound
Y-NHCH2C.tplbond.CR3 [R3 = H,aliphatic, carbocyclic, heterocyclic group;

N-containing heterocyclyl having a H2O-solubilizing group as a

ACCESSION NUMBER: DOCUMENT NUMBER: TITLE: 1995:602373 HCAPLUS

133:183496
Photographic sensitivity increasing alkynylamine compounds and photographic elements
Lok, Roger; Preddy, Carl R.; Harder, John W. Eastman Kodak Co., USA INVENTOR(S): PATENT ASSIGNEE(S):

SOURCE: U.S., 10 pp. CODEN: USXXAM

DOCUMENT TYPE: Patent

English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE A A2 A1 US 5413905 A 19950509 JP 07199390 A2 19950804 EP 665461 A1 19950802 R: BE, CH, DE, FR, GB, IT, LI, NL PRIORITY APPLN. INFO.: US 1993-169833 JP 1994-310508 EP 1994-119839 19931216 19941214 US 1993-169833 A 19931216

OTHER SOURCE(S): MARPAT 123:18325 IT 167307-87-1 167307-90-6 167307-91-7 167307-92-8 167307-93-9 167307-94-0 167307-96-2 MARPAT 123:183296

167307-96-2
RL: MOA (Modifier or additive use); USES (Uses)
(Photog. sensitivity increasing alkynylamine compde.)
167307-87-1 HCAPLUS
6-Benzothiazolecarboxylic acid, 2-(2-butynylamino)-, monosodium salt (SCI)

(CA INDEX NAME)

• Na

167307-90-6 HCAPLUS 5-Benzothiazolecarboxylic acid, 2-[(3-phenyl-2-propynyl)amino]- (9CI)

INDEX NAME)

ANSWER 14 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

• •

167307-96-2 HCAPLUS 2-Butyne-1-sulfinic acid, 4-(2-benzothiazolylamino)-, monopotassium salt (9CI) (CA INDEX NAME)

ANSMER 15 OF 19 HCAPLUS COPYRIGHT 2006 ACS ON STN Entered STN: 17 Mar 1995 A photog. element is described comprising a Ag halide emulsion, the emulsion comprising Ag halide grains which contain an alkynylamine The presence of the alkynylamine dopant imparts to the photog, element advantageous characteristic of increased sensitivity without requiring addition of oxidents to control fog.
ACCESSION NUMEER: 1995:420770 HCAPLUS
DOCUMENT NUMBER: 122:201130
TITLE: Photograph: Photographic elements containing alkynylamine dopants Preddy, Carl R.; Lam, Wai K.; Lok, Roger Eastman Kodak Co., USA PATENT ASSIGNEE (S): SOURCE: U.S., 9 pp. CODEN: USXXAM DOCUMENT TYPE: Patent English PAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE US 5389510 JP 07199391 A A2 B2 19950214 US 1993-169832 JP 1994-313437 19931216 19950804 JP 3440152 PRIORITY APPLN. INFO.: 20030825 US 1993-169832 A 19931216 R SOURCE(S): MARPAT 122:201130 85902-43-8 161765-65-7 161765-68-0 161765-70-4 161765-71-5 RL: MOA (Modifier or additive use); USES (Uses) (dopant for photog. emulsions) 85902-43-8 HCAPLUS 2-Benzothiazolamine, N-2-propynyl- (9CI) (CA INDEX NAME) OTHER SOURCE(S): ин− сн2− с== сн 161765-65-7 HCAPLUS
2-Benzothiazolamine, N-2-butynyl- (9CI) (CA INDEX NAME) мн-сн₂-с==с-ме 161765-68-0 HCAPLUS
2-Benzothiazolamine, N-(3-phenyl-2-propynyl)- (9CI) (CA INDEX NAME) ANSWER 16 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN Entered STN: 25 Nov 1984 NR2CHC≡CR3 AB The fungicidal title compds. I [R = H, Cl-4 alkyl, CO2H, CHO, Cl-4 hydroxyalkyl, mono- and dialkylaminomethyl, Cl-4 alkoxycarbonyl, hydroxyiminomethyl, (un) substituted carbamoyl, Ph; Rl = H, Cl-4 alkyl, Cl-4 alkoxycarbonyl, halo; RRl = (un) substituted benzo; R2 = H, Cl-4 alkyl; R3 = H, iodo) were prepared Thus 2-(methylamino)-5-methylhiazole was treated with BuLi and HC.tplbond.CCH2Br to give 73% I (R = R3 = H, Rl = R2 = Me) (II), which was treated with BuLi and iodine to give 83% I (R = H, Rl = H, Rl = R2 = Me, R3 = iodo). At 900 ppm II completely controlled C s.
1994:591886 HCAPLUS
101:191886
Propynylaminothiazole derivatives
Makisumi, Yesuo; Murabayashi, Akira; Tawara, Katsuya;
Watanabe, Yoshihachi; Takahaehi, Toshio
Shionogi and Co., Ltd., Japan
Eur. Pat. Appl., 42 pp.
CODEN: BPXXDW
Patent PATENT ASSIGNEE(S): SOURCE: DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: Patent English PATENT INFORMATION:

PATENT NO.

EP 111904
EP 111904
R: BE CH, DI
JP 59112978
US 4535088
2A 8309037
ES 527795
AU 8232110
AU 565850
DK 8105872
GB 2132617
GB 2132617
CA 1212117
PRIORITY APPLAN. INFO:: DATE KIND APPLICATION NO. DATE DATE
----19840627
19860122
, LI, NL, SE
19840629
19850813
19840725
19851001
19840622
19871001
19840622
19840629
19860529 A2 A3 FR, A2 A A1 A1 B2 19831215 EP 1983-112646 JP 1982-225271 US 1983-557365 ZA 1983-9037 ES 1983-527795 AU 1983-22110 19821221 19831202 19831205 19831206 DK 1983-5872 GB 1983-33982 19831220 19831221 CA 1983-443876 JP 1982-225271 OTHER SOURCE(6): CASREACT 101:191886; MARPAT 101:191886

IT 92677-39-9P 92677-40-2P 92677-53-7P
92677-54-8P 92677-55-0P
RL: BAC (Biological activity or effector, except adverse); BSU
(Biological
study, unclassified); SPN (Synthetic preparation); BIOL (Biological
study); PREP (Preparation)

Young, Shawquia, Page 15

ANSWER 15 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

161765-70-4 HCAPLUS
2-Benzothiazolemine, N-2-butynyl-5-methyl- (9CI) (CA INDEX NAME)

161765-71-5 HCAPLUS 2-Benzothiazolamine, N-2-butynyl-5-chloro- (9CI) (CA INDEX NAME)

L8

ANSWER 16 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) (prepn. and fungicidal activity of) 92677-39-9 HCAPLUS 2.Benzothiazolamine, N-(3-iodo-2-propynyl)-N-methyl- (9CI) (CA INDEX

92677-40-2 HCAPLUS
2-Benzothiazolamine, N-(3-iodo-2-propynyl)- (9CI) (CA INDEX NAME)

92677-53-7 HCAPLUS
2-Benzothiazolamine, N-(3-iodo-2-propynyl)-N,4-dimethyl- (9CI) (CA INDEX NAME)

92677-54-8 HCAPLUS 2-Benzothiazolamine, N-(3-iodo-2-propynyl)-N,5,6-trimethyl- (9CI) (CA INDEX NAME)

2-Benzethiazolamine, N-(3-iodo-2-propynyl)-N,6-dimethyl- (9CI) (CA INDEX NAME)

ANSWER 16 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

92677-56-0 HCAPLUS 2-Benzothiazolamine, N-(3-iodo-2-propynyl)-6-methoxy-N-methyl- (9CI) (CA INDEX NAME)

92677-87-7P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation and hydrolysis of)
92677-87-7 HCAPLUS
Acctamide, N-2-benzothiazolyl-N-2-propynyl- (9CI) (CA INDEX NAME) IT

85902-43-8P 92677-77-5P 92677-90-2P 92677-91-1P 92677-92-4P 92677-93-5P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation and iodination of) 85902-43-8 HCAPLUS 2-Benzothiazolamine, N-2-propynyl- (9CI) (CA INDEX NAME) ΙT

92677-77-5 HCAPLUS

ANSWER 16 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

92677-64-0P 92677-65-1P 92677-66-2P 92677-84-4P 92677-85-5P 92677-86-6P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of) 92677-64-0 HCAPLUS 2-Benzothiazolamine, N-(3-iodo-2-propynyl)-N,4-dimethyl-, monohydrochloride (9CI) (CA INDEX NAME)

● HCl

92677-65-1 HCAPLUS
2-Benzothiazolamine, N-(3-iodo-2-propynyl)-N,5,6-trimethyl-,
monohydrochloride (9CI) (CA INDEX NAME)

• HC1

92677-66-2 HCAPLUS
2-Benzothiazolamine, N-(3-iodo-2-propynyl)-6-methoxy-N-methyl-, ethanedioate (9CI) (CA INDEX NAME)

CRN 92677-56-0 CMP C12 H11 I N2 O S

ANSWER 16 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) 2-Benzothiazolamine, N-methyl-N-2-propynyl- (9CI) (CA INDEX NAME)

92677-90-2 HCAPLUS 2-Benzothiazolamine, N,4-dimethyl-N-2-propynyl- (9CI) (CA INDEX NAME)

92677-91-3 HCAPLUS 2-Benzothiazolamine, N,5,6-trimethyl-N-2-propynyl- (9CI) (CA INDEX NAME)

92677-92-4 HCAPLUS 2-Benzothiazolamine, N,6-dimethyl-N-2-propynyl- (9CI) (CA INDEX NAME)

92677-93-5 HCAPLUS
2-Benzothiazolamine, 6-methoxy-N-methyl-N-2-propynyl- (9CI) (CA INDEX NAME)

ANSWER 16 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

CM 2

92677-84-4 HCAPLUS 2-Benzothiazolamine, N-(3-iodo-2-propynyl)-N-methyl-, mononitrate (9CI) (CA INDEX NAME)

CM 1

CRN 92677-39-9 CMF C11 H9 I N2 S

7697-37-2 H N O3

92677-85-5 HCAPLUS
2-Benzothiazolamine, N-(3-iodo-2-propynyl)-N-methyl-, monohydrochloride
(9C1) (CA INDEX NAME)

ANSWER 16 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

● HC1

92677-86-6 HCAPLUS 2-Benzothiazolamine, N-(3-iodo-2-propynyl)-N-methyl-, monohydrobromide (9CI) (CA INDEX NAME)

• HBr

ANSWER 17 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)
RL: USES (Uses)
(photogo, element contg., for increased speed and reduced latent image fading)
85902-43-8 HCAPLUS
2-Benzothiazolamine, N-2-propynyl- (9CI) (CA INDEX NAME)

ANSWER 17 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN Entered STN: 01 Sep 1984

A photog. latent image stabilizer comprises a compound of the formula I R1 - H, an aromatic nucleus, or together complete a fused aromatic nucleus: R2

H, Me; Z = O, S, Se, or NR3 where R3 = H or C1-5 alkyl; 21 = CH2NR3, NR3).

Photog. elements containing I also exhibit increased speed. Thus, a multilayer color photog. element was prepared which contained 2-IN-(2-propynyl)aminojbenzoxazole (II) at 0.2 mmol/mol Ag in a faster yellow dye-forming emulsion layer. The element was then cut into 3 parts.

One part was imagewise exposed and processed immediately, a 2nd was

KIND	DATE	APPLICATION NO.	DATE
			• • • • • • • •
A	19840529	US 1983-466244	19830214
λ	19830329	US 1981-320794	19811112
A1	19851001	CA 1983-430255	19830613
н	19841204	US 1984-577934	19840207
		US 1981-320794 A	19811112
		US 1983-466244 A	19830214
	A A A1	A 19840529 A 19830329 A1 19851001 H 19841204	A 19840529 US 1983-466244 A 19830329 US 1981-320794 A1 19851001 CA 1983-430255 H 19841204 US 1984-57934 US 1984-320794 A2

IT 85902-43-8

ANSWER 18 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN Entered STN: 12 May 1984 For diagram(s), see printed CA Issue. A photog, emulsion which exhibits increased speed and reduced latent

image fading contains as a latent image stabilizer a compound of the formula I

= O, S, Se, NR; Y = the necessary atoms to complete a fused aromatic

- O, S, Se, NR; Y - the necessay atoms to complete a fused aromatic nucleus;

Z - CH2NR, NR; and R - H, Cl-5 alkyl). Thus, a multileyer color film element, containing in a faster yellow dye-forming emulsion layer a S-Au sensitized Agbr (1.62 g Ag/m2, gelatin 1.72 g/m2) emulsion, a yellow dye-forming coupler 0.33 g/m2 and II 0.2 emol/mol Ag, was imagewise exposed, stored 2 wk at 25.6 (relative humidity 501), and processed to give an image with a relative blue speed of 170 vs. 83 for a II-free control.

ACCESSION NUMBER:

DS0.CUMENT NUMBER:

99:13920

Photographic speed-increasing and latent image-stabilizing compounds, silver halide emulsions,

99:13920
Photographic speed-increasing and latent
image-stabilizing compounds, silver halide emulsions,
and photographic elements
Lok, Roger; Freeman, John P.; Baum, William N.
Eastman Kodak Co., USA

INVENTOR(S): PATENT ASSIGNEE(S): SOURCE:

U.S., 7 pp. CODEN: USXXAM Patent DOCUMENT TYPE: LANGUAGE:

English 2 PAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 4378426	A	19830329	US 1981-320794	19811112
US 103803	н	19840103	US 1982-408536	19820816
CA 1173042	A1	19840821	CA 1982-411798	19820921
JP 58090634	A2	19830530	JP 1982-196880	19821111
JP 59042293	B4	19841013		
US 4451557	A	19840529	US 1983-466244	19830214
US 104903	н	19841204	US 1984-577934	19840207
PRIORITY APPLN. INFO.:			US 1981-320794 A	3 19811112

US 1983-466244

A3 19830214

85902-43-8 RL: USES (Uses)

(photog, color material containing, for increased speed and latent image

stabilization)

85902-43-8 HCAPLUS 2-Benzothiazolamine, N-2-propynyl- (9CI) (CA INDEX NAME)

```
L8
ED
GI
AB
                 CHBrCH2Br), or COR3 (where R3 = CH2F, OCH2CH:CH2, SEt, OEt, SMe, or
                useful as insecticides, acaricides, or nematocides, were prepared a) by reaction of 2-amino(or alkylamino)benzothiazoles or -benzoxazoles with PCH2COC1 or with PCH2CO2Na and SOC12 or PC13 or b) by reaction of 2-(chloroacetamido)benzothiazoles or -benzoxazoles with KP, followed
2-(cnioroscetamido)benzothiazoles or -benzoxazoles with KF, followed
(when
R1 = H) by treatment with NaH and R1Br or with Me3COK and C1COR3.
Pesticidal compns. containing 1 were reported.

ACCESSION NUMBER: 1972:552162 HCAPUUS
DOCUMENT NUMBER: 77:152162
TITLE: 2-(Pluoroscetamido)benzothiazoles and -benzoxazoles
INVENTOR(S): Bader, Joerg
ATENTA SSIGNEE(S): Agripat S. A.
SOURCE: GET. Offen., 44 pp.
CODEN: GWXXBX
DOCUMENT TYPE: Patent
LANGUAGE: German
PAMILY ACC. NUM. COUNT: 1
```

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 2206575	A	19720914	DE 1972-2206575	19720211
NL 7202462	A	19720829	NL 1972-2462	19720224
DD 100622	С	19731005	DD 1972-161108	19720224
BE 779837	A1	19720825	BE 1972-114359	19720225
FR 2127807	A5	19721013	PR 1972-6518	19720225
ZA 7201255	A	19721129	ZA 1972-1255	19720225
IT 953462	A	19730810	IT 1972-21070	19720225
ES 400151	A1	19751116	ES 1972-400151	19720225
PRIORITY APPLN. INFO.:			CH 1971-2897 A	19710226
			CH 1971-16121 A	19711105

CH 1972-1225

A 19720126

37968-18-6P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of) 37968-18-6 HCAPLUS Acetamide, N-2-benzothiazolyl-2-fluoro-N-2-propynyl- (9CI) (CA INDEX NAME)

L8 ANSWER 19 OF 19 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

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☐ SKEWED/SLANTED IMAGES
☐ COLOR OR BLACK AND WHITE PHOTOGRAPHS
GRAY SCALE DOCUMENTS
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